PATENT COOPERATION TREATY

PCT

REC'D 1/6 DEC 2004

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's	or agent's file reference	FOR FURTHER ACTION		of Transmittal of International	
3157		FOR FURTHER ACTION	Preliminary Exa	mination Report (Form PCT/IPEA/416)	
	al application No. 02/00045	International filling date (day/mon 22.08.2002	th/year)	Priority date (day/month/year) 22.08.2002	
Internation	al Patent Classification (IPC) or t	ooth national classification and IPC			
A61K47/					
Applicant					
Applicant PAPAIO	ANNOU, Dionysios et al.				
1. This	international preliminary exa nority and is transmitted to th	amination report has been prepa e applicant according to Article 3	red by this Intel 86.	rnational Preliminary Examining	
2. This	This REPORT consists of a total of 5 sheets, including this cover sheet.				
×	This report is also accompanied by ANNEXES, i.e. sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT).				
The	These annexes consist of a total of 4 sheets.				
3. This	s report contains indications r	relating to the following items:			
1	☑ Basis of the opinion				
ii	☐ Priority				
111	•	f opinion with regard to novelty,	nventive step a	nd industrial applicability	
IV	☐ Lack of unity of Inver				
V	☐ Reasoned statement		rd to novelty, in	ventive step or industrial applicability;	
Vi	☐ Certain documents c	ited .			
VII	☐ Certain defects in the	international application			
VIII	☐ Certain observations	on the international application			
Date of submission of the demand			Date of completion of this report		
03.07.2003			2.2004		
Name and mailing address of the international preliminary examining authority:			ized Officer	"Reduction Palaticopy.	
- 3	European Patent Office D-80298 Munich	Lüde	mann, S		
9)	Tel. +49 89 2399 - 0 Tx: 523 Fax: +49 89 2399 - 4465	8656 epmu d	none No. +49 89 2	2399-7842	

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/GR 02/00045

1. E	Basis	of the	report
------	-------	--------	--------

Description, Pages

1. With regard to the **elements** of the international application (Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17)):

	1-23	3	as origin	ally filed	
	Clai	ms, Numbers			
	2-7		received	on 03.11.2004 with letter of 03.11.2004	
	1		received	on 23.11.2004 with letter of 23.11.2004	
	Dra	wings, Sheets			
	1/12	-12/12	as origir	nally filed	•
2.	With lang	n regard to the langua luage in which the inte	ge, all the elemernational applic	ents marked above were available or furnished ation was filed, unless otherwise indicated unde	to this Authority in the r this item.
	The	se elements were ava	ilable or furnish	ed to this Authority in the following language:	, which is:
		the language of a trai	nslation furnishe	ed for the purposes of the international search (u	ınder Rule 23.1(b)).
		the language of public	cation of the inte	ernational application (under Rule 48.3(b)).	
		the language of a train Rule 55.2 and/or 55.3	nslation furnishe 3).	ed for the purposes of international preliminary e	xamination (under
3.	With	n regard to any nucle o rnational preliminary e	otide and/or am examination was	nino acid sequence disclosed in the internation carried out on the basis of the sequence listing	al application, the :
		contained in the inter	national applica	tion in written form.	
		filed together with the	e international a	pplication in computer readable form.	
		furnished subsequen	tly to this Autho	rity in written form.	
		furnished subsequen	tly to this Autho	rity in computer readable form.	
	□ ·	The statement that the in the international ap	ne subsequently oplication as file	furnished written sequence listing does not go l d has been furnished.	peyond the disclosure
		The statement that the listing has been furnished	ne information re shed.	ecorded in computer readable form is identical to	the written sequence
4.	The	amendments have re	esulted in the ca	incellation of:	
		the description,	pages:		
	×	the claims,	Nos.:	8-11	
		the drawings,	sheets:		

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/GR 02/00045

5.	This report has been established as if (some of) the amendments had not been made, since they have been considered to go beyond the disclosure as filed (Rule 70.2(c)).
	USELL COUSIDERED TO DEVOLOCULO GOLOGO CO MOSTA CALLANTA

(Any replacement sheet containing such amendments must be referred to under item 1 and annexed to this report.)

6. Additional observations, if necessary:

V. Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

Novelty (N) Yes: Claims 1-7

No: Claims -

Inventive step (IS) Yes: Claims 1-7

No: Claims -

Industrial applicability (IA) Yes: Claims 1-7

No: Claims -

2. Citations and explanations

see separate sheet

Re Item V

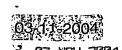
Reasoned statement with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

- 1.1 Reference is made to the following documents:
 - D1: WO 98/34646 A (ATTERWILL CHRISTOPHER KENNETH ;PURCELL WENDY MARIA (GB); ISMAIL FY) 13 August 1998 (1998-08-13)
 - D2: MANFREDINI STEFANO ET AL: "Retinoic acid conjugates as potential antitumor agents: Synthesis and biological activity of conjugates with Ara-A, Ara-C, 3(2H)-furanone, and aniline mustard moieties." JOURNAL OF MEDICINAL CHEMISTRY, vol. 40, no. 23, 7 November 1997 (1997-11-07), pages 3851-3857, XP002236863 ISSN: 0022-2623
 - D3: US-B-6 344 2061 (GIACOMONI PAOLO ET AL) 5 February 2002 (2002-02-05)
 - D4: KARIGIANNIS GEORGE ET AL: "Structure, biological activity and synthesis of polyamine analogues and conjugates." EUROPEAN JOURNAL OF ORGANIC CHEMISTRY, 2000, pages 1841-1863, XP002236864
 - D5: PAPADIMOU EVANGELIA ET AL: "Inhibition of ribonuclease P activity by retinoids." JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 273, no. 38, pages 24375-24378, XP002237316 ISSN: 0021-9258
- 1.2 D1 (WO9834646), which is considered as the closest prior art, discloses antioxidants, e.g. carotene-like substances like retinoic acid linked to targeting moiety such as polyamines, like spermine and spermidine for the treatment of neurodegenerative disorders. Conjugates of polyamines with all-trans-retinoic acids analogues with the structures as disclosed in present claim 1 are not disclosed.
- 1.3 D2 (XP002236863) discloses diamine linked to retinoid disclosed for the treatment of tumors. Substances according to claim 1 are not disclosed.
- 1.4 In D3 (US6344206B1), composition comprising retinol and a polyamine polymer are disclosed. Substances according to claim 1 are not disclosed.
- 1.5 D4 (XP002236864) is a review dealing with polyamine analogues and conjugates.

EXAMINATION REPORT - SEPARATE SHEET

4

- Substances according to claim 1 are not disclosed.
- 1.6 D5 (XP002237316) discloses the inhibition of ribonuclease P activity by retinoids. Substances according to claim 1 are not disclosed.
- 1.7 None of the documents D1-D5 discloses the **all-trans**-retinoic acids analogues with the structures as disclosed in present claim 1.
- 1.8 Furthermore, D1 does not provide any example of how to prepare conjugates of retinoic acids with spermine or spermidine. The examples provided describe synthesis of conjugates via a one-pot reaction of a benzopyran-type antioxidant with a benzylic-type bromine atom used to alkylate the alpha-amino function of an α,ω-diaminoalkane. The present method differs from D1 in that the conjugates are obtained by succinimidyl esters of all-trans-retinoic acids and consecutive purification by flash column chromatography. This is not disclosed or suggested by any of the documents D1-D5.
- 1.9 Therefore, claims 1-7 fulfill the requirements of Art. 33(2) and 33(3) PCT.



5

11:56



AMENDED CLAIMS

FROM PATRINOS & KILIMIRIS

1. Conjugates of polyamines with acidic retinoids and in particular polyamine amides in which the R group of the acyl group(s) RCO is one of the retinoid residues R1-R6 pointed out in the following pharmaceutically important acidic retinoids and polyene chain-shortened all-trans-retinoic acid analogues :

and said polyamines are:

a) Linear tri-, tetra- and hexa-amines, which conjugates have the following general formulae:

wherein n is 1 to 9

b) conformationally restricted polyamines, which conjugates have the following general formulae:

10

15

20



23-NOV-2004 14:05 FROM PATRINOS & KILIMIRIS

COR HOW HAVE HOUSE TOR HOUSE TOR HOUSE TOR HOUSE TOR HOUSE TO THE HOUSE TOR HOUSE TO HOUSE TOR HOUSE TO HOUSE TOR HOUSE TO HOUSE TOR HOU

c) cyclic polyamines, which conjugates have the following general formulae:

d) branched (dimeric) polyamines, which conjugates have the following general formula:

wherein

R' is COR or (CH2)3NHCOR and R" is COR or (CH2)3NHCOR and n is one of the numbers 1, 2 and 7

P. 13

- 2. A method for the preparation of a compound according to claim 1 involving either the following two steps:
 - a) synthesis of compounds with the general formula

5

wherein R is one of the retinoid residues R¹-R⁵ of claim 1, which involves esterification of acidic retinoids with HOSu in the presence of the coupling agent DCC and purification with flash column chromatography b) direct selective acylation of the primary amino groups of polyamines with the as above obtained compounds, or the acylation of the secondary amino groups of polyamines, protected at their primary amino functions with the trifluoroacetyl or the 9-fluorenylmethoxycarbonyl group, with the acidic retinoids of claim 1 in the presence of the coupling agent PyBrOP, followed by deprotection.

15

20

10

3. A method according to claim 2, which method involves the direct selective acylation of the primary amino functions of polyamines or their corresponding hydrochloride or trifluoroacetate salts with the compounds of step a) of claim 2, wherein the solvent is selected between dichloromethane, chloroform and dimethylformamide and the base, where necessary, is selected between triethylamine and diisopropylethylamine or any other tertiary amine or in general any other non-nucleophilic base.

25

4. A method according to claim 3 characterized in that the selective acylation of the primary amino functions of polyamines is effected with any other activated carboxylic acid derivative known to acylate selectively primary amino functions in the presence of secondary ones. 5

10

15

11:56



- 5. A method according to claim 2 characterized in that the selective mono- or bisacylation of primary amino functions of polyamines takes place indirectly and involves the following steps:
 - (i) protection of the secondary amino functions of polyamines, bearing the trityl protecting group at their primary amino functions, with the 9-fluorenylmethoxycarbonyl or the trifluoroacetyl group
 - (ii) detritylation
 - (iii) mono- or bis-acylation with the compounds of step a) of claim 2
 - (iv) complete deprotection and purification, if necessary, by flash column chromatography.
- 6. A method according to claim 2 characterized in that the selective acylation of the secondary amino functions of polyamines involves the following steps:
 - (i) selective trifluoroacetylation of the primary amino functions of polyamines
 - (ii) acylation of the secondary amino functions with the acidic retinoids of claim 1 in the presence of the coupling agent PyBroP
 - (iii) removal of the trifluoroacetyl groups by alkaline hydrolysis.
- 7. Pharmaceutical preparations or products containing the compounds claimed in claim 1 for therapeutical applications in humans